Functional characterization and physiological roles of the single Shaker outward $K^+$ channel in Medicago truncatula

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To cite this version:

Alice Drain, Julien Thouin, Limin Wang, Martin Boeglin, Nicolas Pauly, et al.. Functional characterization and physiological roles of the single Shaker outward $K^+$ channel in Medicago truncatula. Plant Journal, Wiley, 2020, 102 (6), pp.1249-1265. 10.1111/tpj.14697. hal-02457197

HAL Id: hal-02457197
https://hal.archives-ouvertes.fr/hal-02457197

Submitted on 19 Nov 2020
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Article type : Original Article

Title
Functional characterization and physiological roles of the single Shaker outward K⁺ channel in
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Running title
The single outward Shaker channel in Medicago

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This article has been accepted for publication and undergone full peer review but has not been through the copyediting, typesetting, pagination and proofreading process, which may lead to differences between this version and the Version of Record. Please cite this article as doi: 10.1111/tpj.14697

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SUMMARY

The model legume *Medicago truncatula* possesses a single outward Shaker K⁺ channel, while *Arabidopsis thaliana* possesses two channels of this type, named AtSKOR and AtGORK, the former having been shown to play a major role in K⁺ secretion into the xylem sap in the root vasculature and the latter to mediate the efflux of K⁺ across the guard cell membrane leading to stomatal closure. Here we show that the expression pattern of the single *M. truncatula* outward Shaker channel, which has been named MtGORK, includes the root vasculature, guard cells and root hairs. As shown by patch-clamp experiments on root hair protoplasts, besides the Shaker-type slowly-activating outwardly-rectifying K⁺ conductance encoded by MtGORK, a second K⁺-permeable conductance, displaying fast activation and weak rectification, can be expressed by *M. truncatula*. A KO mutation resulting in absence of MtGORK activity is shown to weakly reduce K⁺ translocation to shoots, and only in plants engaged in rhizobial symbiosis, but to strongly affect the control of stomatal aperture and transpirational water loss. In legumes, the early electrical signaling pathway triggered by Nod factor perception is known to comprise a short transient depolarization of the root hair plasma membrane. In absence of MtGORK functional expression, the rate of the membrane repolarization is found to be decreased by about 2 times. This defect was without any consequence on infection thread development and nodule production in plants grown *in vitro*, but a decrease in nodule production was observed in plants grown in soil.

Keywords

*Medicago truncatula*, outward Shaker channels, K⁺ transport to shoots, guard cells, transpiration control, root hairs, Nod factor induced electrical signal
INTRODUCTION

Potassium (K⁺) can compose up to 10% of the total plant dry weight. This major inorganic constituent of the living cell is the most abundant cation in the cytosol, where it is involved in various functions such as electrical neutralization of negatively charged molecules and control of cell membrane polarization. As an unbound highly mobile abundant osmolyte, K⁺ is also involved in regulation of the cell osmotic potential and related functions such as cell growth or osmotically driven cell and organ movements. It also plays a role in the activation of enzymes, protein synthesis, cell metabolism, and photosynthesis (Clarkson and Hanson, 1980, Nieves-Cordones et al., 2016). Thus, plant growth requires that large amounts of K⁺ ions are taken up by roots from the soil solution and distributed throughout the plant. Several tens of membrane transport systems, which belong to at least 3 families of channels, named Shaker, TPK/KCO and TPC, and 3 families of transporters, named HAK, HKT and CPA, contribute to K⁺ transport (uptake, distribution and compartmentalization) in plants (Mäser et al., 2001; Véry et al., 2014). Among them, the Shaker channel family is the best characterized.

Shaker channels give rise to the main K⁺ conductance of the plasma membrane in most plant cell types (Véry and Sentenac, 2003; Hedrich, 2012). Like their counterparts in animal cells, plant Shaker channels have a tetrameric structure, associating 4 Shaker polypeptides, called alpha-subunits (Daram et al., 1997). A Shaker alpha-subunit consists of a hydrophobic core displaying six transmembrane segments, named S1 to S6, and a pore loop, named P, present between S5 and S6 and carrying the hallmark motif GYGD that plays a central role in the channel selectivity for K⁺. The assembly of the four S5-P-S6 modules in the center of the tetrameric protein structures the K⁺ permeation pathway. Plant Shaker channels, like animal Shakers, are regulated by voltage. The S4 segment harbors positively charged residues (H, R and K) and constitutes the channel voltage sensor. The cytosolic C-terminal part, which begins just after the end of S6, displays a C-linker domain, a cyclic-nucleotide binding domain (CNBD), an ankyrin domain (absent in some alpha-subunits), and a KHA domain rich in hydrophobic and acidic residues (Daram et al., 1997; Nieves-Cordones et al., 2014; see supplementary Figure S1b).

The plant Shaker channel family is strongly conserved, each plant genome harboring about 10 Shaker genes that can systematically be sorted into 5 groups, based on phylogenetic and functional analyses (Véry et al., 2014; see supplementary Figure S1a). Group 1 and 2 members (5 members in Arabidopsis) are characterized as inwardly rectifying channels, mediating K⁺ uptake across the cell membrane. Group 3 channels (a single member of this type in Arabidopsis) display a weak rectification and can thus contribute to both K⁺ uptake and secretion across the cell membrane. Group 4 comprises also a single member in Arabidopsis. It
is considered a regulatory subunit since it seems unable to form functional channels by itself but can interact with alpha-subunits from groups 1, 2, and 3 to form heteromeric inward channels with modulated functional features. The last group, group 5, gathers outwardly rectifying channels dedicated to K$^+$ secretion from the cell. It comprises 2 members in Arabidopsis, AtSKOR and AtGORK. AtSKOR is strongly expressed in the root vasculature where it plays a major role in K$^+$ secretion into the xylem sap and thereby in K$^+$ translocation from roots to shoot (Gaymard et al., 1998). AtGORK expression has been detected in various tissues and cell types, including guard cells and root hairs. In guard cells, AtGORK has been shown to encode the outward conductance that mediate the efflux of K$^+$ leading to reduced guard cell turgor allowing stomatal closure (Ache et al., 2000; Hosy et al., 2003). In root hairs, the physiological role of AtGORK is still unclear. In root periphery cells, AtGORK has been shown to mediate an efflux of K$^+$ upon the depolarization of the cell membrane that resulted from a strong increase in the external concentration of Na$^+$ (Shabala and Cuin, 2007). It has been suggested that, by mediating outward K$^+$ currents, AtGORK could contribute to membrane repolarization and thereby prevent detrimental depolarization in response to abiotic or biotic stresses/signals. A role in electrical signaling of elicitors has also been proposed (Ivashikina et al., 2001). Similarly, in legumes, it has been suggested that outwardly-directed K$^+$ currents mediated by AtGORK homologs could play a role in electrical signaling triggered by Nod-factor perception (Damiani et al., 2016a).

So far, besides the work on AtSKOR and AtGORK, very few studies have been aimed at characterizing the functional properties of plant Shaker channels from group 5, and no reverse genetics analysis has highlighted the roles of these functionally-characterized channels (Langer et al., 2002; Sano et al., 2007; Huang et al., 2018). Here we investigate the functional properties and roles of MtGORK, the unique member of the Shaker group 5 in the model legume Medicago truncatula.

RESULTS

Molecular cloning and primary structure of MtGORK
Phylogenetics analyses indicate that M. truncatula Shaker channel group 5 comprises a single member (Damiani et al., 2016a; Wang et al., 2019), Medtr5g077770, hereafter named MtGORK (Figure S1a). The corresponding cDNA (2508 bp) was amplified by PCR, allowing sequence analysis of the deduced polypeptide (Figures S1b and S1c) and determination of the gene structure (Figure S1d). MtGORK possesses the Shaker channel typical hydrophobic core, with the 6 transmembrane segments S1-S6 and the pore loop harboring the GYGD hallmark motif between S5 and S6 (Figure S1b and S1c). A C-linker domain, a cyclic nucleotide binding domain,
an ankyrin domain and a K$_{HA}$ domain can be identified in the C-terminal region downstream the hydrophobic core, like in the Arabidopsis AtSKOR and AtGORK outward channels. The percentages of identity and similarity between MtGORK and AtSKOR or AtGORK are close to 70% and 85%, respectively. The two residues of the P loop and 2 residues of the S6 transmembrane segment that have been identified in AtSKOR as contributing to the dependency of the channel voltage-sensitive gating on the external concentration of K$^+$ (Johansson et al., 2006), conserved in AtGORK, are also present in MtGORK (Figure S1c).

Functional characterization in Xenopus oocytes
Depolarization of the membrane elicited an outward current in oocytes injected with MtGORK cRNA and not in control oocytes injected with water (Figure 1a). The exogenous macroscopic current displayed slow sigmoidal activation kinetics and reached a steady-state value within ca. 2 s (Figure 1a). Steady-state I-V curves displayed a strong outward rectification (Figure 1b). Comparison of the I-V curves obtained in presence of 10, 30 or 100 mM K$^+$ in the external solution revealed that increasing this concentration resulted in a positive shift of the activation potential threshold, i.e., the threshold beyond which, when the potential was shifted to more positive values, outward currents became detectable (Figure 1b). The experimental curves describing the dependency of the channel relative open probability, $P_o/P_{o\text{max}}$, on voltage (obtained from MtGORK deactivation currents recorded at +50 mV after pre-pulses varying from -100 mV to +80 mV) in presence of 10, 30 or 100 mM external K$^+$ were fitted with the classical two-state Boltzmann law (Ache et al., 2000). The results indicated that the channel half activation potential ($E_{a50}$: membrane potential at which the channel relative open probability is 0.5) was strongly dependent on external K$^+$ concentration, being shifted by +50 mV when this concentration was increased from 10 to 100 mM (Figure 1c). Such a regulation by external K$^+$ ensures that the outward rectification of MtGORK is total regardless of the concentration of K$^+$ prevailing outside, and thus that this channel is strictly dedicated to K$^+$ release. Besides the sensitivity to voltage and external K$^+$, MtGORK was also found to be sensitive to the external pH, the outward current being decreased by about 50% when the pH was decreased from 7.5 to 5.6 (Figure 1d). Thus, the pH sensitivity of MtGORK appears stronger than that reported in AtGORK (Ache et al., 2000).

A last series of experiments was aimed at characterizing the ionic selectivity of MtGORK. As expected, the current reversal potential, $E_{rev}$, determined from classical analysis of tail current recordings (Ache et al., 2000), was found to be dependent on the external concentration of K$^+$ (Figure 1e). $E_{rev}$ shifted by about +50 mV for a 10-fold increase in external K$^+$ concentration, indicating that MtGORK displays a strong selectivity for K$^+$ (under several theoretical
assumptions, a channel exclusively permeable to K\(^+\) is predicted to give rise to a shift of ca. +58 mV upon a ten-fold change in K\(^+\) concentration; Hille, 2001). Shifts in \(E_{\text{rev}}\) were also recorded upon replacement of K\(^+\) in the external medium by another alkali cation, either Li\(^+\), Na\(^+\), Rb\(^+\) or Cs\(^+\), at the same concentration (100 mM) (Figure 1f). In such experiments, the magnitude of the \(E_{\text{rev}}\) shift reflects the relative permeability of the substituting cation and allows to determine this permeability, reported to that to K\(^+\), using the so-called Goldman equation (Hille, 2001). MtGORK displayed the following permeability sequence (Eisenman's series IV; Eisenman, 1961), K\(^+\)>Rb\(^+\)>Cs\(^+\)>Na\(^+\)=Li\(^+\) (Figure 1f), which is identical to that reported for AtSKOR (Gaymard et al., 1998). Like in AtSKOR, the relative permeability to Rb\(^+\) is rather high (ca. 0.6), while that to Na\(^+\) is weak (<0.1).

**A KO mutation in MtGORK results in absence of Shaker type outward K\(^+\) conductance in guard cells**

A *M. truncatula* line (cv Jemalong A17) named NF9352, displaying an insertion of the Tnt-1 retrotransposon in the first exon of MtGORK (Figure S1d) was obtained from the mutant collection of the Noble Foundation and self-pollinated to produce the F2 generation. The plant that was amplified possessed this mutation, hereafter named mtgork, at the hemizygous state. Genotyping experiments (PCR) were carried out to identify both mutant plants homozygous for the mtgork mutation and control WT plants possessing a wild type (WT) genotype at this locus. RT-PCR experiments could not amplify bona fide MtGORK transcripts in mutant plants homozygous for the mtgork mutation (Figure 2a), providing evidence that mtgork is a knock-out (KO) mutation. The selected mutant and WT lines were further amplified for phenotyping experiments. Visual observations of the plants homozygous mutant or WT for the mtgork mutation did not allow to detect any specific phenotype. Furthermore, measurements of root and shoot biomass production in plants grown in different conditions (*in vitro*, in greenhouse or growth chamber on compost or sand-vermiculite mixture) at different growth stages and inoculated with a rhizobial strain (*S. meliloti* 1021 strain) or not inoculated did not reveal any specific defect in plant development (Figure S2).

Patch-clamp experiments carried out on guard cell protoplasts revealed that homozygous plants for the mtgork mutation did not display the typical Shaker-like outward K\(^+\) currents that were recorded in WT plants (Figure 2b). Small instantaneously-activating outward K\(^+\) currents were detected in the mutant protoplasts (Figure 2b and 2c). On the other hand, the inward K\(^+\) currents were very similar in mutant and WT protoplasts. Based on their activation kinetics and current-voltage curve, these currents are likely to be mediated by inwardly rectifying Shaker channels like in Arabidopsis guard cells (Lebaudy et al., 2008).
The *mtgork* KO mutation results in impaired control of transpirational water loss

A sharp reduction in the rate of leaf transpirational water loss rapidly occurred (<20 min) in WT plants after leaf excision, whereas no significant change in this rate could be observed in the mutant plants over 60 min (Figure 3a). The dotted lines plotted in Figure 3a correspond to data obtained in a similar experiment in Arabidopsis, with mutant plants that did not express the *AtGORK* gene shown to encode the guard cell outward Shaker conductance (Hosy et al., 2003).

Figure 3b shows that impaired control of leaf transpirational water loss in *mtgork* mutant leaves rapidly resulted, within less than 60 min after leaf excision, in a strong withered phenotype. A high rate of transpirational water loss still occurred at that time in the mutant leaves, while WT leaves appeared at the same time to have almost maintained their initial "turgid" phenotype and displayed no longer significant transpirational loss (Figure 3a and 3b).

In a parallel experiment, excised leaves were periodically photographed using a macroscope allowing to observe stomata and to compare the kinetics of stomatal closure in the two genotypes (Figure 4). The photographs displayed by Figure 4a show that, in WT leaves, the stomata appeared as already tightly close 10 min after leaf excision, when compared with their initial state observed just after leaf excision. Indeed, the kinetics of stomatal closure obtained from the analysis of such photographs shows a very rapid reduction, within 10 min, in stomatal aperture in WT leaves (Figure 4b). In *mtgork* mutant leaves, in contrast, the stomatal aperture observed 10 min after leaf excision appeared poorly different from that observed just after leaf excision (Figure 4a). The kinetics of stomatal closure in the mutant leaves displayed by Figure 4b confirmed that stomatal closure was strongly impaired in the mutant leaves.

**MtGORK** is expressed in root stellar tissues and root hairs

**MtGORK** expression was first investigated *in silico* using public databases ["eFP Browser Medicago", (http://bar.utoronto.ca/), and "Medicago Gene Atlas", (http://mtgea.noble.org/v3/)]. The data indicated that **MtGORK** is expressed in leaves, stem, roots and nodules. Then, analysis of *M. truncatula* roots transformed with a **MtGORK** promoter-GUS reporter gene construct indicated that the expression pattern of **MtGORK** includes root hairs, root vascular tissues and nodule vasculature (Figure 5). Evidence for the expression of **MtGORK** in root hairs has also been provided by RNA-Seq analyses (Damiani et al., 2016a).

In Arabidopsis, the outward Shaker **AtGORK** is expressed in root hairs and **AtSKOR** in the root stele. We have investigated the consequences of the *mtgork* mutation in these root tissues/cell types through the analysis of (i) the translocation of K⁺ towards the shoots, (ii) the
electrical properties of the root hair cell membrane, and (iii) the early electrical signal induced by Nod factor (NF) perception in root hairs and the plant capacity to engage rhizobial symbiosis.

**K⁺ translocation towards the shoots**
Experiments carried out to assess the contribution of MtGORK to K⁺ translocation to shoots revealed that the mtgork mutation poorly affected the shoot K⁺ content. A reduction in this content could be observed in inoculated plants, but it was slight (ca. 13%). No statistically significant difference was observed in non-inoculated plants (Figure 6). In agreement with these results, the absence of MtGORK activity was found to be without any significant effect on the concentration of K⁺ in exuded xylem sap and on the volume of exuded sap (due to root pressure) upon shoot excision, whatever the plant status, inoculated or non-inoculated (Figure 7).

**Electrical properties of the root hair plasma membrane**
Patch-clamp recordings in 28 root hair protoplasts from WT plants revealed that two distinct types of outward K⁺ conductances (Figure 8a; upper panels), differing at least in their activation kinetics and in the shape and size of the deactivation current, as previously reported (Wang et al., 2019), could dominate the membrane permeability to K⁺. Here, in experiments made in a native context, we use the term "conductance" to refer to a type of permeation pathway that can be mediated by a single molecular identity or by a set of channels of several identities but displaying similar properties.

In some protoplasts (10 out of 28), the dominant conductance displayed a slow sigmoidal activation of currents and slow current deactivation kinetics upon return to the holding voltage (Figure 8a, left panel), reminiscent of the features of MtGORK when expressed in Xenopus oocytes (Figure 1c) or when characterized in situ at the guard cell membrane in WT plants (Figure 2b, left panel), and typical of outward Shaker channels (Gaymard et al., 1998; Ache et al., 2000; Langer et al., 2002; Sano et al., 2007; Huang et al., 2018). Such a conductance has also been recorded in root hairs from Arabidopsis (Ivashikina et al., 2001) and Medicago sativa (Bouteau et al., 1999). On the other hand, this type of conductance was not observed in mutant plants homozygous for the mtgork mutation (Figure 8a, lower panel).

The second conductance that could be identified as the dominant one in WT protoplasts (in 13 out of 28 protoplasts) was also found in mtgork mutant root hair protoplasts (Figure 8a, upper central panel and lower panel, respectively). The recorded current traces upon membrane depolarization reveal an instantaneous component and a time-dependent component, the relative part of the latter increasing with depolarization (Figure 8a). The fact that the time-dependent component of this second conductance was also displayed by the current traces obtained in
mutant protoplasts that did not express MtGORK, indicates that this conductance does not result from the addition of a MtGORK component to an instantaneously-activating conductance. Previous analyses of this conductance (in another *M. truncatula* genotype) have revealed that it is less selective for K$^+$ than the Shaker-type one and that it displays a weak apparent voltage gating (apparent gating charge $z = 0.2$; Wang et al., 2019). This conductance was thus qualified as a weakly voltage-gated outwardly rectifying cationic conductance, mostly already open at the time the positive voltage is imposed. Another characteristic of this second conductance is that the deactivation currents observed upon return to the holding voltage are small when compared with those of outward Shaker channels like MtGORK. This conductance was named fast-activating outward cationic conductance by Wang et al. (2019) since it displays a more rapid activation than that of the Shaker-type outward conductance. The name "fast-activating" conductance is also used in the present report.

In a few WT protoplasts (5 out of 28), the recorded traces indicated that the two above described conductances were simultaneously active at the root hair cell membrane: *e.g.*, a large instantaneously activating component could be distinguished together with a slowly activating component and large and slowly deactivating tail currents (Figure 8a, upper right panel).

In summary, the outward conductance of the membrane to K$^+$ could be considered in most WT protoplasts (23 out of 28) as essentially resulting from the activity of a single conductance type, either the fast-activating conductance or the MtGORK one, these two situations having rather similar frequencies (the former conductance being the dominant one in 13 protoplasts and the MtGORK conductance the dominant one in 10 protoplasts, out of the 28 as indicated above). Sorting the 28 protoplasts into these two categories and deriving the corresponding current-voltage curves revealed that, in contrast to the protoplasts dominated by the fast-activating conductance, the protoplasts dominated by the MtGORK conductance displayed a strong outward rectification (Figure 8b), reminiscent of that displayed by oocytes expressing MtGORK (Figure 1b). Finally, the current-voltage curve derived for this set of 28 WT protoplasts, treated as a whole, was similar, in terms of current magnitude, to the one obtained for 15 protoplasts from mutant plants homozygous for the *mtgork* mutation (Figure 8c). Altogether, these results suggest that the absence of MtGORK conductance in root hairs of the mutant plant was compensated by an increase in the expression or activity of the fast-activating outward conductance. It is worth to note that the patch-clamp recordings carried out in guard cell protoplasts from WT and *mtgork* mutant guard cell protoplasts (Figure 2) did not reveal any conductance likely to correspond to the root hair fast activating conductance.
MtGORK contributes to repolarization of the root hair cell membrane following Nod Factor induced depolarization

Initiation of symbiotic interactions with N\textsubscript{2}-fixing rhizobia in legumes is triggered at the root hair cell membrane in response to nodulation factors (NF) secreted by rhizobia, and involves complex signaling events (Felle et al., 1998; Oldroyd and Downie, 2008). The earliest events that have been reported, together with ROS production (Puppo et al., 2013; Damiani et al., 2016b), are changes in ion fluxes, H\textsuperscript{+}, Ca\textsuperscript{2+}, anion (Cl\textsuperscript{-}) and K\textsuperscript{+}, through the root hair plasma membrane resulting in a transient depolarization of this membrane (Felle et al., 1998). Continuous recordings of the local concentrations of Ca\textsuperscript{2+}, H\textsuperscript{+}, Cl\textsuperscript{-} and K\textsuperscript{+} at the root surface using extracellular ion selective micro-electrodes and of the changes in membrane potential using an intracellular micro-electrode have shown that NF perception rapidly results (within ca. 1 min) in an increase in net Ca\textsuperscript{2+} influx, followed by a net efflux of anions and possibly an inhibition H\textsuperscript{+}-excretion. Altogether, these events result in a strong membrane depolarization, which activates voltage-sensitive K\textsuperscript{+} channels, allowing an efflux of K\textsuperscript{+} that repolarizes the membrane, a process to which an activation (or a re-activation) of H\textsuperscript{+} excretion by proton pumps could contribute (Felle et al., 1998). The hypothesis that MtGORK contributes to the efflux of K\textsuperscript{+} involved in the repolarization of the membrane during this "action potential-like" signaling process was tested by comparing the kinetics of membrane repolarization, recorded by microelectrode impalement, in WT and mutant plants as described in Figure 9. The membrane potential before NF treatment was not significantly different between WT and mutant cells (-138 ± 14 mV and -141 ± 20 mV, respectively; n=5). The NF-induced depolarization was found to be larger, although not statistically different, in WT than in mutant cells, -34 ± 11 and -25 ± 6 mV (means ± SD; n=5), respectively. Interestingly, the rate of repolarization (mV.min\textsuperscript{-1}) was found to be significantly different between the two types of plants, about two times slower in mutant plants homozygous for the mtgork disruption when compared with the control WT plants (Figure 9c).

MtGORK activity is not necessary for infection thread development and nodule formation

The number of infection threads developed in plants grown \textit{in vitro} on Farhæus medium and observed at either 3 or 5 days after root inoculation (dpi) with \textit{S. meliloti} was not significantly different between mutant plants homozygous for the mtgork disruption and control WT plants (Figure 10a). In agreement with this result, when growth continuously occurred \textit{in vitro} on Farhæus medium, the number of nodules determined at either 14 or 21 dpi was similar in the mutant and WT plants (Figure 10b). In contrast, experiments carried out using plants grown in soil for 25 days revealed a larger number of nodules in WT than in mutant plants (Figure 10c).
DISCUSSION

A single outwardly rectifying Shaker channel in *M. truncatula*

The Shaker channel family comprises a single outwardly rectifying member in *M. truncatula* (Supplemental Figure S1a). It is interesting to note that *in silico* analysis of the genome sequence of *Lotus japonicus* (https://lotus.au.dk/) reveals a single outwardly rectifying member in the Shaker family of this legume model too. In Arabidopsis, the Shaker gene family comprises 2 genes coding for outwardly rectifying channels, *AtGORK* and *AtSKOR*, and 7 genes for inwardly (or weakly-inwardly) rectifying channels (Supplemental Figure S1a). The phylogenetic structure of the Shaker family is strongly conserved in plants: about 10 genes and always 5 subgroups, the genes coding for inwardly-rectifying channels (groups 1 to 4) being more numerous than those coding for outwardly-rectifying channels (group 5) (Véry *et al.*, 2014).

Analyses of the functional properties of plant inwardly rectifying channels in heterologous systems have revealed distinctive features and regulations, in terms of voltage sensitivity, affinity for external K⁺ or sensitivity to pH (Véry and Sentenac, 2003; Véry *et al.*, 2014). Formation of heteromeric channels associating subunits encoded by different inward Shaker genes can further increase this functional diversity (Reintanz *et al.*, 2002; Duby *et al.*, 2009; Jeanguenin *et al.*, 2011). The corresponding information presently available on outwardly rectifying channels is much more reduced. Besides MtGORK and the Arabidopsis *AtGORK* (Ache *et al.*, 2000; Hosy *et al.*, 2003) and *AtSKOR* (Gaymard *et al.*, 1998; Johansson *et al.*, 2006), only 3 other outward K⁺ channels have been characterized at the functional levels in heterologous systems, NTORK from tobacco (Sano *et al.*, 2007), PTORK from poplar (Langer *et al.*, 2002) and CmSKOR from melon (Huang *et al.*, 2018). These outward Shaker channels share a common functional feature: their activation threshold depends on the external concentration of K⁺. This regulation by external K⁺, also observed in outward K⁺ conductances recorded *in planta* (Schroeder, 1989; Blatt, 1991; Blatt and Gradmann, 1997; Roelfsema and Prins, 1997; Wang *et al.*, 2019), ensures that the channels open only when the transmembrane K⁺ electrochemical gradient is outwardly directed, regardless of the external concentration of K⁺, and thus that these channels are dedicated to the function of K⁺ secretion into the apoplasm/external medium. Dominated by this tight regulation, the functional diversity of outward Shaker channels in plants seems to be rather reduced, when compared to that of inward Shaker channels. This might explain that the number of outward Shaker genes is low in every plant genome.
Both MtGORK and a non-Shaker K⁺-permeable outward conductance can be expressed in root hairs

Differences in the current activation kinetics and in the shape and size of deactivation currents displayed by protoplasts from wild-type plants supported the hypothesis that two types of K⁺-permeable outward conductance could be active at their plasma membrane. These two types of conductance do not correspond to two different cell types, e.g. trichoblasts versus atrichoblasts or cortical cells, since they have also been observed in spheroplasts obtained from the tip of young elongating root hairs using a laser-mediated procedure allowing root hair selection (Wang et al., 2019). Finally, the fact that M. truncatula root hairs can express two distinct types of K⁺-permeable outward conductance is further evidenced by the finding that one of these conductances can be also observed in protoplasts from root hairs of mutant plants that do not express MtGORK.

In summary, one of these two conductances is dependent on MtGORK functional expression and displays the same functional features as MtGORK when heterologously expressed in oocytes. It thus results from MtGORK activity. The second one, named fast-activating outward cationic conductance as in Wang et al. (2019), cannot be hypothesized to result from the activity of another Shaker channel. Indeed, within the M. truncatula Shaker family, MtGORK is the only outward channel gene. All the other genes belong to groups 1 to 4 (Supplemental Figure S1) and are thus likely to code for inwardly- or a weakly inwardly-rectifying channels that display activation upon membrane hyperpolarization. A simple hypothesis, based on the present knowledge, is that this fast-activating outward conductance corresponds to either a Cyclic-Nucleotide-Gated Channel (CNGC), a Glutamate Receptor (GLR) or an Annexin. The electrophysiological properties of the members from these families are still poorly characterized (Hedrich, 2012).

This K⁺-permeable fast-activating outward conductance is reminiscent of conductances reported in other cell types and plant species, for instance the NORC conductance characterized in barley root xylem parenchyma (Wegner and Raschke, 1994; Wegner and De Boer, 1997) and the weakly voltage-dependent non-selective cation conductance described in wheat roots (Davenport and Tester, 2000). However, this type of conductance was not reported in previous electrophysiological analyses carried out in root hairs from Arabidopsis (Ache et al., 2000, Ivashikina et al., 2001) and alfalfa (Bouteau et al., 1999; Kurkdjian et al., 2000). This might indicate that the levels of expression or activity of this conductance in root hairs is strongly dependent on the plant material, growth conditions and plant species, or is specific of M. truncatula when compared with Arabidopsis and alfalfa.

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In most wild-type root hair protoplasts (ca. 80% of the protoplasts), the membrane outward conductance to K⁺ was strongly dominated by either the MtGORK or the fast-activating conductance, with rather similar frequencies, and current features revealing that the two conductances were simultaneously active could be detected in only ca. 20% of the protoplasts. Hence, it is likely that heterogeneities in the electrical properties of the root hair plasma membrane exist, at least transiently, amongst young root hairs. Such heterogeneities might be related to those of root hair tip responses to Nod Factors (NF), since NF treatment do not result in an alteration of tip growth in the same way in every young root hair (Esseling et al., 2004).

Comparison of the roles of MtGORK and AtGORK in control of stomatal aperture

In guard cells, the membrane outward K⁺-permeable conductance was dominated by MtGORK (Figure 2). It is interesting to note that no fast-activating outward conductance similar to that expressed by root hairs was detected in guard cells (Figure 2b), and that the absence of MtGORK activity in the mutant plants did not appear to be compensated by a conductance of this type (Figure 2b and 2c).

In Arabidopsis guard cells, AtGORK dominates the membrane outward conductance to K⁺ and mediates the depolarization-induced potassium release involved in stomatal closure (Ache et al., 2001). Mutant plants harboring a KO mutation in AtGORK displayed slower closure kinetics, when compared with wild-type plants, resulting in impaired control of transpirational water loss (Hosy et al., 2003). The data shown by Figures 3 and 4 reveal that the role of AtGORK in Arabidopsis guard cell, allowing efficient stomatal closure, is played by MtGORK in M. truncatula. Furthermore, they suggest that the control of leaf transpirational water loss and the contribution of the Shaker outward conductance to this function are stronger in M. truncatula than in Arabidopsis (dotted lines in Figure 3a; withered phenotype within less than 60 min in Figure 3b).

Comparison of the roles of MtGORK and AtSKOR in K⁺ translocation to shoots

The mtgork KO mutation was found to be without any significant effect on the shoot K⁺ content in non-inoculated plants (Figure 6a), and to result in a slight reduction of this content in plants inoculated with the symbiotic partner S. meliloti (Figure 6b). Also, the concentration of K⁺ in the xylem sap driven by root pressure (after shoot excision) and the volume of exuded sap were not significantly affected by the mtgork mutation (Figure 7). Thus, absence of outward Shaker channel activity in M. truncatula root stele poorly affects K⁺ translocation towards the shoots, especially when compared with the corresponding results reported in Arabidopsis, where the absence of outward Shaker channel activity due to a KO mutation in AtSKOR results in a reduction in shoot K⁺ content and K⁺ concentration in the xylem sap by about 50% (Gaymard et
This indicates that MtGORK poorly contributes to K⁺ translocation towards the shoots, in contrast to AtSKOR, or that its absence in the mutant plants is efficiently compensated by other types of K⁺-permeable conductances. Non-Shaker K⁺-permeable outward conductances have been identified in xylem parenchyma cells from barley (Wegner and Raschke, 1994; Wegner and De Boer, 1997).

The K⁺ concentration of the collected xylem sap samples was about 5 times higher in the inoculated than non-inoculated plants (Figure 7). However, the shoot K⁺ contents were larger only by about 1.5 times in the former than in the latter plants (Figure 6). Furthermore, in the same experimental conditions, the shoot biomass was lower by about 2 times in the former than in the latter plants (Supplemented Figure S2a and S2b). This suggests that the flux of recirculated K⁺ ions from shoots to roots via the phloem sap is larger in symbiotic conditions. A larger flux of phloem sap towards the roots would provide sugars to functioning nodules. MtGORK, which is expressed in the nodule vasculature (Figure 5), may thus contribute to the recirculation towards the shoots of K⁺ ions arriving in nodules via the phloem sap. This hypothesis is consistent with the fact that, when inoculated, WT plants displayed (slightly) higher shoot K⁺ contents than mutant plants (Figure 6).

Role of MtGORK in root hairs and early transduction of Nod Factor signal

In Arabidopsis root hairs, AtGORK encodes the typical K⁺-sensitive voltage-gated outwardly-rectifying conductance (Ache et al., 2000), without any significant contribution of AtSKOR (Ivashikina et al., 2001). AtGORK has been hypothesized in this cell type to be involved in control of cell turgor and membrane potential. In connection with this latter function, it has also been hypothesized to play a role in electrical signals (transient depolarization and changes in K⁺ fluxes) such as those induced by elicitor treatments (Ivashikina et al., 2001). It should however be noted that none of these hypotheses has received support from reverse genetics approaches so far, and thus that the role of AtGORK in Arabidopsis root hairs is still unclear.

Here we show that MtGORK contributes to the repolarization of the root hair cell membrane following the NF induced depolarization (Figure 9). The repolarization still occurred in absence of MtGORK activity, but at a slower rate, by about two times (Figure 9c). Thus, these results indicate that MtGORK plays a role in the electrical signal triggered by NF perception by contributing to the membrane repolarization. It is however not the only electrogenic transport system involved in this repolarization process since repolarization is observed, although at a slower rate, in mutant root hairs devoid of MtGORK conductance. The simplest hypothesis is that the fast-activating outward conductance identified in root hairs both in presence or absence of
MtGORK functional expression (Figure 8) plays a role in the repolarization and can compensate for the absence of MtGORK activity in mutant plants.

We checked whether the slower repolarization of the root hair cell membrane in the mutant plants affected subsequent steps of plant engagement in the symbiotic interaction with S. meliloti. Comparison of the number of infection threads in WT and mutant plants grown in vitro on agar plates in presence of S. meliloti did not reveal any significant consequence of the mtgork mutation (Figure 10a). Nodule production was also similar in the two types of plants when grown in vitro (Figure 10b). These results indicate that the kinetics of membrane repolarization is not a crucial component of the signaling pathway leading to the symbiotic interaction of M. truncatula with S. meliloti when plants are grown on agar plates. The reduction in nodule production by the mtgork mutant, when compared with WT controls, when plants are grown in soil (Figure 10c) might however indicate that plant capacity to engage rhizobial symbiosis in more physiological conditions requires integrity of the early electrical signaling events, but it cannot be excluded that this reduction in nodule production results from more general physiological defects.

**Conclusion**

The most striking difference between MtGORK, representative of all the plant outwardly-rectifying Shaker channels characterized so far, and the fast-activating K⁺-permeable outward conductance, appears to be the rectification capacity. The fast-activating conductance is weakly rectifying and thus can allow K⁺ influx when the electrochemical gradient of this cation across the plasma membrane is inwardly directed, while the strong regulation of MtGORK by both the voltage and the external concentration of K⁺ ensures that the permeation pathway remains closed when the K⁺ electrochemical gradient is inwardly directed, so that these channels are strictly dedicated to K⁺ secretion. Based on the present knowledge, all plant species whose genome has been sequenced possess at least one Shaker channel of this type (Véry et al., 2014). While some species possess 4 genes encoding such channels, like grapevine or poplar (Véry et al., 2014), a single one is sufficient in other species. It is interesting to note that, in species displaying a single outward Shaker channel like M. truncatula, this channel can display a rather broad expression pattern, suggesting that it might be involved in the various functions involving its different orthologs in species that harbor several channels of this type. Within the framework of this hypothesis, the fact that the absence of MtGORK channel activity poorly affects K⁺ translocation towards the shoots, when compared with the effects of the corresponding mutation in Arabidopsis, would result from compensation in some tissues of the absence of MtGORK by other types of conductances in the mutant plants. It should be noted that, in root hairs in contrast
to guard cells, the fast-activating K⁺-permeable conductance appears to be able to compensate the absence of MtGORK conductance in mutant plants. Despite such possibilities of redundancy and compensation, the fact that all plant species possess at least one outward Shaker channel gene indicates that K⁺ channels displaying a strict outward rectification provide important services in some environmental conditions.

EXPERIMENTAL PROCEDURES

Plant material and plant growth

*M. truncatula* (ecotype Jemalong A17) seeds were scarified with sulfuric acid (99%) for 10 min, rinsed and sterilized in 6% sodium hypochlorite solution for 3 min. After 3 h imbibition in sterile water, seed coats were removed and seeds were transferred onto 1% agar plates in Petri dishes, which were turned upside down (agar up) and remained in the dark at 4°C for at least 48 hours in order to break dormancy and obtain synchronization of germination. The plates were then transferred at 21°C for 16 to 24 h for germination. The radicles were then about 2 cm long.

For *in vitro* culture, germinated seedlings were transferred onto a sterile sheet (12 × 8.5 cm) of chromatography paper (Rogo-Sampaic, France) laid on solid Fahraeus agar medium (modified from Vincent, 1970) in a Petri dish (12 × 12 cm, for 10 seedlings). The medium contained 10 g L⁻¹ of purified agar (Euromedex, https://web.euromedex.com/) and 0.5 mM MgSO₄, 0.7 mM KH₂PO₄, 0.8 mM Na₂HPO₄, 1 mM CaCl₂, 20 μM Fe-citrate and 0.1 mg.L⁻¹ of MnSO₄, CuSO₄, ZnSO₄, H₃BO₃ and Na₂MoO₄, pH 7.5 (adjusted with KOH). The lower part of the plate was wrapped in aluminum foil to avoid detrimental effects of light on roots. The plate was placed in a quasi-vertical position in a growth chamber (70% relative humidity, 70 μE.m⁻².s⁻¹ light intensity) with a photoperiod of 16 h light (25°C) and 8 h dark (21°C) for 5 days.

Germinated seedlings grown on the agar plates for 2 further days were transferred on sand-vermiculite mixture (3:1, v:v; ca. 1 L per plant in 10 L containers) or sand-compost (3:1, v:v; ca. 1.5 L per plant in individual pots) and grown in a growth chamber (70% relative humidity, 16 h light, 300 μE.m⁻².s⁻¹ light intensity, 25°C, and 8 h night, 21°C) or in greenhouse, respectively. They were watered twice a week, alternatively with water or Fahraeus solution.

Composite plants were generated according to the protocol of Boisson-Dernier *et al.* (2001) using the electrocompetent *Agrobacterium rhizogenes* strain ARqua1 harboring the transcriptional GUS construct (2 kb *gork* promoter sequence cloned in the pGWB3 vector from Gateway system) (Nakagawa *et al.*, 2007).
Mutant and control wild-type plants

*M. truncatula* mutant line NF9352 was identified (BLASTn of the GORK genomic sequence in the Noble website) in the Noble collection (http://medicago-mutant.noble.org/mutant/) as harboring a Tnt-1 retrotransposon insertion in the MtGORK gene (Supplemental Figure S1d). We named the mutation resulting from this insertion mtgork. A F1 plant hemizygous for the mtgork mutation was amplified. PCR genotyping experiments on the F2 progeny identified plants either homozygous for the mtgork mutation or displaying a wild type genotype at this locus and thereafter named control WT plants. Both types of plants were amplified for F3 progeny.

Rhizobial strain and plant inoculation

The rhizobial strains used for *M. truncatula* inoculation were *Sinorhizobium meliloti* Rm1021, Rm1021 DsRed and Rm2011 LacZ. Bacteria were grown in 5 g.L\(^{-1}\) Bacto tryptone, 3 g.L\(^{-1}\) yeast extract, 6 mM CaCl\(_2\), pH 7.2 (TY/Ca medium), supplemented with the appropriate antibiotic: 50 µg.mL\(^{-1}\) streptomycin for Rm1021 and Rm1021 DsRed and 10 µg.mL\(^{-1}\) tetracycline, centrifuged, resuspended and washed in Fahraeus medium. Aliquots of the final bacterial suspension (OD\(_{600}\) ~ 10\(^{-2}\)) were directly laid over apices of the germinated seedlings. Sand-vermiculite and sand-compost mixtures were also inoculated (ca. 10 mL of the rhizobial suspension for 1 L of soil).

Plant K\(^+\) contents and K\(^+\) translocation to shoots via the xylem sap

Plants were grown on sand-vermiculite mixture for 8 weeks in growth chamber. Shoots and roots were collected, dried and weighted (DW). Ions were extracted with 0.1 N HCl and assayed (flame spectrophotometry). The shoots of plants grown in parallel, in the same conditions and for the same time, were excised below the first leaf, at about 1 cm above the soil. For each plant, about 30 min after shoot excision, the root extremity was introduced into a plastic tube (Eppendorf type) through a hole pierced at the tube bottom and sealed with silicon paste to collect exuded sap for 24 h. The volume and K\(^+\) and Ca\(^{2+}\) concentrations of the exuded sap were determined (weight measurement and flame spectrophotometry assay, respectively).

Two-electrode voltage clamp characterization of MtGORK in Xenopus oocytes

The coding sequence of MtGORK was amplified by PCR, cloned into the pGEM-Xho vector (derived from pGEMDG; D. Becker, Würzburg) downstream from the T7 promoter and between the 5’- and 3’-untranslated regions of the Xenopus β-globin gene. Capped and polyadenylated copy RNA (cRNA) were synthesized *in vitro* (mMESSAGE mMACHINE T7 kit, Ambion). Oocytes, isolated and handled as described previously (Véry *et al.*, 1995), were injected with ca. 30 ng of MtGORK cRNA (ca. 1 ng.nL\(^{-1}\)) or with 30 nl of diethyl-pyrocarbonate-(DEPC) treated water for
control ("water injected") oocytes. They were then kept at 18 °C in ND96 solution (96 mM NaCl, 2 mM KCl, 1.8 mM CaCl₂, 1 mM MgCl₂, 2.5 mM sodium pyruvate, and 5 mM HEPES-NaOH, pH 7.4) supplemented with 0.5 mg L⁻¹ of gentamycin until voltage-clamp recordings. Whole-oocyte currents were recorded using a two-electrode voltage-clamp technique 1–2 d after cRNA injection. All electrodes were filled with 3 M KCl. The external solution bathing the oocyte was continuously percolated during the voltage-clamp experiment. All bath solutions contained a background of 1 mM CaCl₂ and 2 mM MgCl₂, buffered with either 10 mM Heps-NaOH or 10 mM Mes-NaOH, pH 7.5 or 5.6, respectively. This background solution was supplemented with either 100, 30, 10 or 3 mM KCl, the osmolarity of the solution being maintained constant by addition of N-Methyl-D-glucamine (NMDG), or with 100 mM CsCl, RbCl, NaCl or LiCl. To extract MtGORK-mediated currents from total oocyte currents, mean currents recorded in water-injected control oocytes from the same batch in the same ionic conditions were subtracted from those recorded in MtGORK-expressing oocytes.

Leaf transpirational water loss and stomatal aperture measurement

Plants of WT and mtgork lines were grown in greenhouse in compost for 6 weeks. Then, leaves were excised and exposed to desiccation by placing them on trays on the lab bench at room temperature. Leaves were weighed (FW: fresh weight) at different time points. Transpirational water loss was determined from the difference between the weight at each time point and the initial weight. For stomatal aperture measurement, the excised leaves were periodically photographed (magnification: 250x) using a microscope (AxioZoom V16, Zeiss, fitted with a PlanNeoFluar 2.3X objective; https://www.zeiss.fr/microscopie/home.html), equipped with an ORCAFlash4 Hamamatsu camera (https://www.hamamatsu.com/eu/en/index.html).

Protoplast isolation and patch-clamp analyses

Root hair protoplasts were obtained by enzymatic digestion as previously described (Wang et al., 2019) and stored in ice until patch-clamp measurements. For guard cell protoplast preparation, epidermal strips were peeled off from the abaxial surface of 6 to 7 leaves using forceps, and cut into small pieces. The enzymatic treatment was performed for 1h and 40 min at 28°C in a solution containing 1% (w/v) cellulase RS, 0.1% (w/v) pectolyase Y23, 1% (w/v) BSA, 1 mM CaCl₂, 2 mM ascorbic acid, 1 mM Mes, and 450 mM D-mannitol, its pH being adjusted to 5.7 with KOH. Then, the released protoplasts were collected by filtering through a 40 µm nylon mesh and washed with the conservation medium twice and stored in ice until patch-clamp measurements. The conservation medium contained 1 mM CaCl₂, 2 mM ascorbic acid, 1 mM Mes, and 500 mM D-mannitol, its pH being adjusted to 5.7 with KOH.
Patch-clamp experiments were performed in the whole-cell configuration. Patch-clamp pipettes were pulled using a DMZ-Universal Puller (Zeitz-Instruments GmbH, Germany) from borosilicate capillaries (GC150F-7.5; Phymep, France) and fire polished (by the DMZ-Universal Puller). Microelectrode resistance was about 10 and 14 MΩms for patch clamping root hair and, respectively, guard cell protoplasts. A reference Ag/AgCl half-cell completed the circuit. The patch clamp amplifier was an Axopatch 200B (Axon Instruments Inc., USA). Whole-cell currents were measured at least 5 minutes after seal formation. Data were sampled at 1 kHz. The Clampex module of the pClamp9 software (Axon Instruments Inc., USA) was used for data acquisition. Analysis was performed using the Clampfit module of pClamp10 and SigmaPlot 11 (Systat Software Inc., USA). Liquid junction potentials were corrected.

Membrane potential measurements

*M. truncatula* seedlings were grown on agar medium (10 g L⁻¹ of purified agar in a solution, named bath background solution, bbs, containing 0.1 mM KCl, 0.1 mM CaCl₂, 0.1 mM NaCl and 2 mM MES-Bis-Tris-Propane, pH 6.5) for 2 days. Roots were excised and fixed in a plexiglass chamber filled with bbs. The chamber was percolated with bbs for 15 min (recovery treatment) before root impalement. Impalement micro-electrodes, with a tip diameter of approximately 0.5 μm, were pulled from borosilicate glass capillaries (GC200F-10, Harvard Apparatus, http://www.harvardapparatus.com) and back-filled with 3 M KCl. Microelectrodes were connected via an Ag/AgCl pellet to an HS-2 · 0.1L probe of an Axoprobe 1A electrometer (Axon Instruments). The reference comprised a combined glass pH electrode (filled with 3 M KCl) placed in the chamber downstream of the root. The micro-electrode was placed at the root surface, at about 0.5 cm from the tip, in front of young developing root hairs, using a manually operated micro-manipulator (Narishige, http://narishige-group.com). Subsequently, the vertical position of the root chamber was adjusted using a micro-elevator (IT6D CA1, Microcontrole, http://www.newport.com), allowing precise penetration of the micro-electrode into an epidermal cell at 10-40 μm below the root surface. During impalement, the bath solution was continuously refreshed. The steady-state membrane potential was successively measured in 5 external solutions: bbs, bbs + 10⁻⁶ M Nod factor, bbs, bb + 30 mM KCl and bbs again. The whole protocol was achieved within less than 30 min. The recording was discarded when the membrane potential values got in the 3 bbs successively perfused during the protocol were not consistent together.

Promoter fusion and histochemical localization of GUS Activity
A 2-kb DNA fragment upstream of the starting ATG of MtGORK gene (Medtr5g077770), was amplified by PCR using gene-specific primers (pGORK2kb-F1: CACTCCTTAGCAAAATAGCAAAAATTA and pGORK2kb-R1: GAAATTAATATACCTCATCCTTAGAAG). Composite M. truncatula plants were obtained by transformation with Agrobacterium rhizogenes Arqual strain as previously described (Andrio et al., 2013). Healthy composite plants were transferred onto new plates containing modified Fahraeus medium without nitrogen and kanamycin. Plants were inoculated with S. meliloti 3 d after transfer.

Transgenic roots were stained with GUS assay buffer as previously described (Andrio et al., 2013). Roots from at least 20 plants from three biological experiments were examined. Roots and nodules were fixed in 1% glutaraldehyde and 2% formaldehyde in 0.05 M phosphate buffer (pH 7), washed, dehydrated, and embedded in Technovit 7100 according to the manufacturer’s instructions. Fifty-micrometer-thick vibroslices were obtained with a HM560V Vibratome (Leica RM 2165) and visualized with an Olympus BH-2 microscope using dark-field optics.

DATA STATEMENT
All relevant data can be found within the manuscript and its supporting materials. Contact for plant material and further information: M.B. (martin.boeglin@inra.fr) and A.-A.V. (very@supagro.inra.fr)
ACKNOWLEDGEMENTS
This work was supported in part by an ANR grant (ANR-11-BSV7-010-02) (to HS and AAV), a
doctoral grant of the Government of Nouvelle Calédonie (to AD) and from the French Ministry of
Research (to JT). We thank Dr. Carine Alcon for help in stomatal aperture measurements at the
Montpellier BPMP PHIV platform (https://www1.montpellier.inra.fr/wp-inra/bpmp/plateformes-et-
plateaux-techniques/phiv/).

CONFLICT OF INTEREST
The authors declare that there is no conflict of interest.

AUTHOR CONTRIBUTIONS
performed research; A.D., J.T., H.S., A.-A.V, M.B., L.W., M.N.-C., I.G. & N.P. analyzed data; H.S.,
A.D., J.T. & A.-A.V. wrote the paper.

SUPPORTING INFORMATION
Additional Supporting Information may be found in the online version of this article.

Figure S1. MtGORK from the Shaker channel family of Medicago truncatula.

Figure S2. Mutant plants homozygous for the mtgork mutation display similar shoot and root
biomass as wild type plants.
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FIGURE LEGENDS

**Figure 1.** Functional characterization of MtGORK in *Xenopus* oocytes

(a) Voltage clamp protocol (top left) and typical currents recorded in control oocytes injected with H$_2$O (bottom left) or injected with *MtGORK* cRNA (right) in 100 mM K$^+$ solution. Voltage-clamp pulses varied from -100 to +80 mV, in increments of 20 mV. Every episode of imposed voltage lasted 3 seconds. The holding potential was -50 mV.

(b) MtGORK current-voltage relationships at different external K$^+$ concentrations: 10, 30 or 100 mM. Mean ± SE, n ≥ 8.

(c) Effect of the membrane voltage and the external K$^+$ concentration on MtGORK open probability. The relative open probability (Po/Pomax) was obtained from the analysis of deactivation currents upon return to the holding voltage after the activation pulse (mean ± SE, n ≥ 6). The solid lines are Boltzmann fits to the mean Po/Pomax values. The mean values (± standard error, n ≥ 6) of the half-activation potential (Ea50) of the MtGORK channel obtained from these fits in the different concentrations of K$^+$ are provided in the inset.

(d) Activation of MtGORK currents by external alkalization. The external solution contained 100 mM K$^+$. The external pH was 5.6 or 7.5. Means ± SE, n = 4.

(e) Variation of MtGORK reversal potential of currents (E$_{rev}$) with the external concentration of K$^+$ (mean ± SE, n ≥ 8). E$_{rev}$ was determined in each solution using a tail-current protocol: After activation of MtGORK channels at +60 mV, voltage pulses were performed at lower voltages flanking Erev, and Erev was obtained from the analysis of the deactivation currents. Dashed line: K$^+$ equilibrium potential (E$_{K^+}$).

(f) Permeability of MtGORK to different monovalent cations. (Left) MtGORK deactivation currents were recorded using a tail-current protocol (as above) in bath solutions containing 100 mmole.l$^{-1}$ of either K$^+$, Rb$^+$, Cs$^+$ Na$^+$ or Li$^+$. Mean deactivation currents ± SE, n≥ 6. (Right) Permeability ratios of the different cations with respect to that of K$^+$, calculated (from the variations of E$_{rev}$) using the Goldman-Hodgkin-Katz equation. Mean ± SE, n≥ 6.

**Figure 2.** The *mtgork* mutation results in absence of MtGORK functional expression in guard cells.

(a) PCR analyses reveal that mutant plants homozygous for the *mtgork* mutation do not express *MtGORK* transcripts. Plants homozygous for the *mtgork* mutation (mutant) or displaying a wild type genotype for this mutation (WT) were grown *in vitro* on Fahraeus medium for 10 days before RNA extraction from whole plants. PCR experiments did not detect *MtGORK* transcripts in the
mutant plants. The gene MtSLAH2/3-b (Medtr6g045200) (Damiani et al., 2016a) was taken as control (lower panel).

(b and c) Comparison of K⁺ currents in guard cell protoplast from WT and mtgork mutant plants. Currents were recorded using the patch-clamp technique in the whole-cell configuration. The ionic composition of the bath solution was 10 mM K-glutamate, 10 mM CaCl₂ and 10 mM Mes-Tris, pH 5.8. Patch-clamp pipette solution: 100 mM K-glutamate, 5 mM EGTA, 1 mM CaCl₂ (free Ca²⁺: 20 nM), 2 mM MgCl₂, 2 mM Mg-ATP, 10 mM Hepes-Tris, pH 7.5. The osmolarity of bath and pipette solutions was adjusted to 480 and 500 mosmol/Kg with D-mannitol. “Outward” whole-cell currents were recorded applying successive pulses of clamped voltage from -71 to 49 mV (after liquid junction potential (LJP) correction) in 15 mV increment from a holding potential at -71 mV. Linward currents were recorded applying voltage pulses from -71 to -161 mV in -15 mV increment from a holding potential at -71 mV. (b): Representative current traces recorded in WT and mutant protoplasts. (c): Steady state current-voltage (I-V) plot. Means ±SE (WT:  n = 5; mutant: n = 9).

Figure 3. The mtgork mutation results in impaired transpirational water loss control. Plants homozygous for the mtgork mutation or displaying a wild type genotype for this mutation were grown on compost in greenhouse for 6 weeks. Leaves were excised during the light period. (a) Water loss was determined by monitoring the decrease in fresh weight of the excised leaves. Means ± SE, n = 3. (b) Photographs of WT and mtgork leaves taken just after excision (t=0) and 60 min after excision (representative of five independent experiments).

Figure 4. The mtgork mutation affects stomatal closure. Leaves from WT and mtgork mutant plants were excised and periodically photographed using a macroscope to compare the kinetics of stomatal closure in the two genotypes. (a) Photographs (representative of at least 3 independent experiments) taken just after excision and, of same leaf zones, 10 min after excision. (b) Kinetics of stomatal closure derived from the analysis of the photographs. Means ± SE (n > 62 stomata from 3 leaves for both genotypes).
**Figure 5.** Expression pattern of the MtGORK gene in roots.

Histochemical analysis of GUS activity in *M. truncatula* composite plants expressing a pMtGORK::GUS reporter construct. MtGORK promoter activity was detected in root epidermis (a), root hairs (b), root stellar tissues (c) and nodules (d).

**Figure 6.** Effect of the absence of MtGORK activity on root and shoot K⁺ contents in non-inoculated and inoculated plants.

Plants homozygous for the *mtgork* mutation (mutant) or displaying a wild type genotype for this mutation (WT) were transferred onto sand-vermiculite mixture 3 days after germination. Shoots and roots were collected for K⁺ and Ca²⁺ assays after 8 weeks of growth. Means ± SE; n = 12.

(a) Non inoculated plants.
(b) Inoculated plants. Inoculation was performed with *S. meliloti* strain 1021 at the end of the second week of growth.

**Figure 7.** Absence of MtGORK activity does not impact the xylem flux of K⁺ from roots to shoot.

Plants homozygous for the *mtgork* mutation (mutant) or displaying a wild type genotype for this mutation (WT) were grown as described in the legend to Figure 6. After 8 weeks of growth, shoots were excised at about 1 cm above the soil. About 30 min after shoot excision, exsuded sap was collected for 24 h. The volume and K⁺ and Ca²⁺ concentrations of the exuded sap were measured. Means ± SE; n = 12.

(a) Non inoculated plants.
(b) Inoculated plants. Inoculation was performed with *S. meliloti* strain 1021 at the end of the second week of growth.

**Figure 8. Outward K⁺ currents in root hair protoplasts from WT or mutant plants.**

Root hair (RH) protoplasts were enzymatically obtained from plants homozygous for the *mtgork* mutation (mutant) or displaying a wild type genotype for this mutation (WT). The bath solution contained 30 mM K-gluconate, 1 mM CaCl₂, 10 mM Mes-Tris, pH 5.6. Patch-clamp pipette solution: 150 mM mM K-gluconate, 10 mM EGTA, 2 mM MgCl₂, 2 mM Mg-ATP, 10 mM Hepes-Tris, pH 7.4. The osmolarity of the bath and pipette solutions was adjusted to 290 and 300 mosmol/Kg respectively, with D-sorbitol. Voltage clamp protocol: pulses from -75 to +45 mV, 15 mV increment, and holding potential at -75 mV.
(a) Representative current traces recorded in WT and mutant protoplasts. In WT protoplasts (top panels), three types of macroscopic conductances could be identified. The top left recording is typical of outward Shaker channels (slow and sigmoidal activation, large deactivation currents). The top middle example can be ascribed to the fast-activating outward cationic conductance previously described (Wang et al., 2019). This conductance displays an instantaneous component and a time-dependent component, and small deactivation currents upon return to the holding voltage. The top right recording provides an example of protoplast where these two types of conductances can be detected as simultaneously active. In mutant protoplasts (lower panel), the outward Shaker conductance was not observed, while the fast-activating cationic outward conductance was present.

(b) Steady state current-voltage (I-V) relationship of the outward Shaker type conductance and of the fast-activating outward cationic conductance derived from patch-clamp recordings in 28 WT protoplasts. The membrane outward conductance to K⁺ was dominated by the outward Shaker type conductance in 10 protoplasts, and by the fast-activating cationic conductance in 13 protoplasts. In 5 protoplasts, a contribution of both conductances to the outward current could be distinguished. I-V curves were derived for the 10 protoplasts dominated by the Shaker-type conductance (squares) and the 13 protoplasts dominated by the fast-activating conductance (diamonds). Means ± SE.

(c) Steady state current-voltage relationship of the membrane outward conductance in WT and mutant protoplasts. Means ± SE. WT: n = 28 (same protoplasts as in panel b, all the protoplasts being taken into account without distinguishing between them and their dominating conductance). Mutant: n = 15.

Figure 9. Role of MtGORK in repolarization of the root hair cell membrane after the initial depolarization induced by Nod Factor perception.

(a) Representative recordings of membrane potential variations induced by NF factor treatment in WT (upper panel) and mtgork mutant (lower panel) plants. The external solution bathing the root (bath background solution: BG) when the micropipette was impaled in an epidermal cell of the root hair zone contained 0.1 mM KCl, 0.1 mM CaCl₂, 0.1 mM NaCl and 2 mM MES-Bis-Tris-Propane, pH 6.5. When a stable signal was observed, the BG solution was replaced by Nod Factor solution (BG + NF: 10⁻⁶ M Nod Factor in BG). The treatment resulted in a rapid depolarization of the cell membrane followed by a repolarization to the initial value while NFs were still present in the percolated solution. Then, the BG + NF solution was replaced by BG, allowing to check that the membrane potential went back close to the initial value recorded before
the NF treatment. The BG was thereafter supplemented with 30 mM KCl (BG + KCl) to check the depolarizing effect of a high K\(^+\) concentration. The 30 mM KCl solution was thereafter replaced by BG to check whether the impaled cell could still restore its membrane potential to the initial value. When the whole test procedure appeared positive, the recording was used for analyzing the repolarization phase during the NF treatment.

(b) Analysis of the repolarization phase. The quasi-linear part of the repolarization phase (lasting about 1 min, from ca. 0.1 min after the beginning of this phase) was extracted and fitted with a linear regression (dashed line) to derive the mean slope (in mV.min\(^{-1}\)) of the recording, which was taken as an estimate of the repolarization rate. Black and grey curves: example of recordings obtained in a plant homozygous for the mtgork mutation (mutant) or displaying a wild type genotype for this mutation (WT), respectively.

(c) Repolarization rates in WT and mutant plants. Means ± SD, n = 5. The asterisk indicates that the difference is statistically significant (Student test, P<0.05).

**Figure 10.** Effect of the mtgork mutation on plant capacity to produce infection threads in presence of rhizobia and on nodule production.

(a) and (b) Germinated seedlings homozygous for the mtgork mutation (mutant) or displaying a wild type genotype for this mutation (WT) were transferred onto Fahraeus agar medium in Petri dishes and inoculated with *S. meliloti* two days later.

(a) Number of infection threads observed at 3 and 5 days post-infection (dpi). The inoculated *S. meliloti* strain was RM2011 lacZ, and the collected roots were stained to reveal LacZ activity for easier detection of infection threads. Means ± SE, n = 10. Statistical analysis (Tukey test at the 5% level) indicates that there was no significant difference between the two types of plants.

(b) Number of nodules at 14 and 21 dpi in plants grown on agar plates. The inoculated *S. meliloti* strain was RM1021 DsRED, and the collected roots were observed using DsRED fluorescence microscopy, allowing easier detection of nodules. Means ± SE, n = 24. No statistically significant difference between the two types of plants (Tukey test at the 5% level).

(c) Nodule production in plants grown in artificial soil. Three days after germination, plants were transferred on artificial soil (sand/compost/clay balls, 2/1/1, v/v/v) and grown in growth chamber. Inoculation (with *S. meliloti* strain RM1021) was achieved 7 days after the transfer. The bacterial suspension prepared for inoculation had an OD close to 0.04. Inoculation was carried out by directly using this suspension or a 1/100 diluted suspension. Roots were collected and gently washed for nodule counting at 18 dpi. Means ± SE; n = 10. Different letters in front of bars indicate that the corresponding difference is statistically significant (Student t-test, P<0.05).
SUPPLEMENTAL FIGURE LEGENDS

Figure S1. MtGORK from the Shaker channel family of *Medicago truncatula*.
(a) Phylogenetic relationships between Shaker polypeptides from *Arabidopsis thaliana* and *Medicago truncatula*. The plant Shaker family comprises 5 groups (see main text). Shaker sequences from Arabidopsis were obtained from the TAIR website (http://www.arabidopsis.org/). A homology search was carried out against the *M. truncatula* protein sequence bank (MT4.0v2) using the BLAST (Basic Local Alignment Search Tool) program, the BLOSUM62 matrix (BLOcks SUBstitution Matrix) and a threshold E (or E-value) equal to 10^{-3}. The unrooted phylogenetic tree was generated with PhyML software (http://www.atgc-montpellier.fr/phyml/binaries.php) using the maximum-likelihood method and 1000 bootstrap replicates in Seaview application (http://doua.prabi.fr/software/seaview). Arabidopsis Shaker polypeptide sequences were first aligned with Muscle (http://www.drive5.com/muscle/), then treated with Gblocks in Seaview program for alignment curation. The phylogenetic tree was drawn with Dendroscope (http://ab.inf.uni-tuebingen.de/software/dendroscope/). Bootstrap values (as percentages) are indicated at the corresponding nodes. The scale bar corresponds to a distance of 10 changes per 100 amino acid positions.

(b) Structure of plant Shaker channels. The channel hydrophobic core comprises 6 transmembrane segments, named S1 to S6. S4 (the so-called voltage sensor) contains positively charged residues and confers sensitivity to the electric field in the membrane (and thus to the transmembrane voltage). P: pore domain. Shaker functional channels are tetrameric proteins, the 4 P domains being assembled in the center of the tetrameric structure where they form the K⁺ permeation pathway (pore). Four large domains can be identified in the cytosolic region downstream S6: a C-linker domain, a cyclic nucleotide binding domain, an ankyrin domain (not present in every plant Shaker channel but present in MtGORK, AtSKOR and AtGORK), and a K⁺/HA domain. Role of these domains in plant Shaker channels: see Daram et al., 1997; Nieves-Cordones et al., 2014).

(c) Sequence alignment of MtGORK and the Arabidopsis AtSKOR Shaker in the P domain and S6. Asterisks denote residues shown to be involved in the channel sensitivity to the external concentration of K⁺ in AtSKOR (Johansson et al., 2006).

(d) Schematic diagram of the MtGORK gene structure indicating the site of insertion of the disrupting TNT1 retrotransposon in the Jemalong A17 *M. truncatula* line named NF9352. Same abbreviations as in panel b. Boxes: exons.
Figure S2. Mutant plants homozygous for the *mtgork* mutation display similar shoot and root biomass as wild type plants. Plants homozygous for the *mt gork* mutation (mutant) or displaying a wild type genotype for this mutation (WT) were compared with respect to biomass production in different conditions.

(a) and (b) Non-inoculated (a) or inoculated (b) plants grown for 8 weeks in growth-chamber on vermiculite-sand mixture. Plants were transferred onto this artificial soil 3 days after germination. When inoculated (B), inoculation (with *S. meliloti* strain 1021) was achieved after 7 days of growth on this soil. Shoot and root were collected for biomass measurements (dry weight: DW) after 8 weeks of growth. Means ± SE; n = 12.

(c) Inoculated plants grown in vitro for 3 weeks. Germinated seedlings were transferred onto Fahraeus agar medium in Petri dishes and inoculated with *S. meliloti* strain 1021 DsRED. Plant dry weight was measured 21 days post-inoculation (means ± SE; n = 24).

(d) Inoculated plants grown for 2, 3 or 4 weeks on compost in growth-chamber. Germinated seedlings were transferred on compost and inoculated one week later with *S. meliloti* strain 1021. Plants were collected for dry weight measurements at 7, 14 and 21 dpi.
Figure 1.
Figure 2.

(a) Western blots showing expression of *MtGORK* and *MtSLAH2/3-b* in WT and mutant guard cell protoplasts.

(b) Current-voltage (I-V) curves for WT and mutant guard cell protoplasts.

(c) Voltage-gated ion channel current traces for WT and mutant guard cell protoplasts.
Figure 3.

(a) Water loss (% initial fresh weight) over time after leaf excision (min).

(b) Comparison of water loss between WT and mutant at different time points.

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Figure 4.

(a) WT and mutant stomatal aperture at t = 0 and t = 10 min.

(b) Graph showing the change in stomatal aperture (µm) over time (min) for WT and mutant samples.
Figure 5.
Figure 6.
Figure 7.
Figure 8.

(a) Outward K⁺ conductances identified

In WT RH protoplast
Shaker type
Fast-activating type

In mutant RH protoplasts
Fast-activating type

(b) (c)

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**Figure 9.**

(a) Wild-Type and mutant membrane potential changes over time with different conditions. 
(b) Linear regression analysis showing the repolarization rate for WT and mutant. 
(c) Graph of repolarization rate comparing WT and mutant.
Figure 10.